1	Assessment of dicarbonyl contributions to secondary organic aerosols over China
2	using RAMS-CMAQ
3	Jialin Li ¹ , Meigen Zhang ^{1, 2, 3,*} , Guiqian Tang ¹ , Yele Sun ^{1, 2, 3} , Fangkun Wu ¹ ,
4	Yongfu Xu ^{1,3}
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6	¹ State Key Laboratory of Atmospheric Boundary Layer Physics and Atmospheric Chemistry,
7	Institute of Atmospheric Physics, Chinese Academy of Sciences, Beijing 100029, China
8	² Center for Excellence in Regional Atmospheric Environment, Institute of Urban Environment,
9	Chinese Academy of Sciences, Xiamen 361021, China
10	³ University of Chinese Academy of Sciences, Beijing 100049, China
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12 13	Abstract The concentration of secondary organic aerosol (SOA) is underestimated in current model studies.
14	Recent research suggests that the aqueous uptake of dicarbonyls contributes to the production of
15	SOA, although few models have included this pathway. Glyoxal, an important representative
16	component of dicarbonyls in models, is significantly underestimated. We therefore incorporated the
17	aqueous uptake of dicarbonyls into the regional air quality modeling system RAMS-CMAQ (the
18	Regional Atmospheric Modeling System-Community Multiscale Air Quality) to evaluate the
19	contribution of dicardonyls to SOA, and we then assess the impact of the underestimation of glyoxal
20	on the production of SOA in China during two time periods: June 3 to July 11, 2014 (episode 1) and
21	October 14 to November 14, 2014 (episode 2). When the aqueous uptake process was added, the
22	modeled mean concentration of SOA in episode 1 increased by 3.65 μ g/m ³ , which explained 34.8%
23	of the unaccounted source of SOA. Whereas the increase in the concentration of SOA in episode 2
24	was 1.82 μ g/m ³ as a result of the lower liquid water content and the lower amount of dicarbonyls
25	produced from biogenic precursors in the fall. On this basis, when the glyoxal simulation was
26	improved, the modeled mean dicarbonyl-derived SOA (AAQ) increased by more than a factor of 2
27	in both episodes relative to case 1. AAQ in episode 1 contributed, on average, 61.0% of the total
28	concentration of SOA and the increase in this contribution represented 70.0% of the unaccounted
29	concentration of SOA, whereas the mean AAQ in episode 2 accounted for 64.5% of total
30	concentration of SOA. Based on the results, the mean AAQ over China was generally higher in the
31	east than in the west during the two episodes. The highest value (10–15 μ g/m ³) of episode 1 appeared

32 in the areas around the lower reaches of the Yellow River. Whereas the highest value of $5-10 \text{ }\mu\text{g/m}^3$ 33 in episode 2 was concentrated over regions from south of the lower reaches of the Yellow River to 34 the south of Guangzhou Province as well as the Sichuan Basin. The contribution of AAQ to the concentration of SOA in episode 1 varied from 10 to 90% throughout China, with the highest 35 contributions (70-90%) in the coastal regions and offshore along the East China Sea to the South 36 37 China Sea and in the southwestern regions. Whereas the fraction of AAQ to SOA in episode 2 was 38 in the range of 10-80% over China, with the fraction up to 80% in a small portion of northeastern 39 China.

40 Keywords: secondary organic aerosol, aqueous uptake, glyoxal, China, RAMS-CAMQ

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42 **1. Introduction**

43 The fine particle fraction of aerosols (PM_{2.5}, i.e., particulate matter with an aerodynamic diameter 44 \leq 2.5 µm) not only absorbs and scatters sunlight in the atmosphere, resulting in a reduction in 45 visibility and effects on the global climate (IPCC, 2007), but also has adverse effects on human 46 health (Harrison and Yin, 2000;Poschl, 2005). Organic aerosol (OA) is a major component of fine 47 particulate matter globally (Murphy et al., 2006;Zhang et al., 2007), typically making up 20-90% 48 of the fine particle fraction (Roberts et al., 2001;Kanakidou et al., 2005;Zhang et al., 2007), 49 suggesting that it has a significant effect on the characteristics and properties of fine particulate 50 matter. Organic aerosols are formed from both primary (direct emissions) and secondary sources. 51 Secondary organic aerosols (SOAs) are formed from precursors of volatile organic compounds 52 (VOCs), which can be either natural or anthropogenic. However, the underestimation of organic 53 aerosol has become a major issue in almost all current atmospheric models due to the incomplete 54 representation of SOA (Heald et al., 2005;Morris et al., 2005;Morris et al., 2006;Goldstein and 55 Galbally, 2007; Yu et al., 2008; Fu et al., 2009; Farina et al., 2010; Jiang et al., 2012; Zhang and Ying, 56 2012; Jo et al., 2013; Lin et al., 2016).

It has been reported that the concentration of SOA in the models is underestimated by one to two orders of magnitude (de Gouw et al., 2005;Volkamer et al., 2006). These results have motivated researchers to investigate why these models are predicting SOA concentrations so poorly. Traditionally, improvements in models have mainly concentrated on the gas-phase and derived heterogeneous formation processes, such as the formation of SOA from aromatic compounds under low- and high-NO_x conditions (Ng et al., 2007), the production of SOA from the oxidation of isoprene and its subsequent growth through acid catalysis (Kroll et al., 2005, 2006;Surratt et al., 2007), the use of volatile bias set method to depict the formation of SOA instead of the two-product model (Lane et al., 2008;Murphy and Pandis, 2009;Han et al., 2016;Lin et al., 2016), and the formation of SOA through the uptake of isoprene epoxides on the surface of aerosols (Ying et al., 2015;Hu et al., 2017). The gap has been closed to some degree through the SOA modeling efforts, but there are still large uncertainties (Tsigaridis et al., 2014).

69 Many studies have suggested that glyoxal and methylglyoxal can form SOAs through chemical 70 reactions in cloud or fog water (e.g. Warneck, 2003;Ervens et al., 2004;Lim and Ziemann, 71 2005;Carlton et al., 2006;Loeffler et al., 2006), or by irreversible uptake on the surface of cloud droplets and aqueous aerosols (e.g. Liggio et al., 2005;Corrigan et al., 2008;Galloway et al., 72 73 2009;Ervens and Volkamer, 2010;Lim et al., 2010), which is probably a significant source of SOA 74 (Ervens et al., 2014;Curry et al., 2018). A few studies (e.g. Carlton et al., 2008;Fu et al., 2009;Carlton 75 et al., 2010;Lin et al., 2012;Li et al., 2013;Lin et al., 2014;Woo and McNeill, 2015) have 76 incorporated aqueous SOA formation pathways into atmospheric models. Several of these studies 77 have shown that chemical reactions in cloud or fog water make negligible contribution to near-78 surface SOA relative to irreversible uptake on the surface of cloud droplets and aqueous aerosols, 79 and that the aqueous formation of SOA cannot completely explain the gaps between the observations 80 and simulations. There are still considerable uncertainties in our knowledge of the formation of SOA. 81 A series of studies (Fu et al., 2008;Myriokefalitakis et al., 2008;Liu et al., 2012;Li et al., 2018) 82 has shown that there is a substantial underestimation in the modeled vertical column densities 83 (VCDs) of glyoxal, but few studies have considered the impact of this underestimation on 84 simulations of SOA concentrations. Xu et al. (2017) and Ervens et al. (2011) showed that the 85 aqueous formation of SOA depends on the liquid water content (LWC), which varies between 86 seasons. Previous studies, such as those of Fu et al. (2009) and Li et al. (2013), have only considered 87 the contribution from SOA derived from the aqueous uptake of dicarbonyls (pathway M) in the 88 summer period or over evergreen areas. In this study, we not only incorporated pathway M into the 89 RAMS-CMAQ (the Regional Atmospheric Modeling System-Community Multiscale Air Quality) 90 modeling system to evaluate the corresponding contribution of dicarbonyls to SOA, but also 91 improved the simulation of glyoxal concentrations by investigating the reasons for its

92 underestimation and assessing its impacts on the concentration of SOA during the two episodes in93 the summer and fall of China.

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95 2. Model and data

96 2.1 Base model description

97 Li et al. (2017a) used the RAMS-CMAQ modeling system to investigate the effects of 98 underestimated aromatic VOC emissions and yields of SOAs from different gas precursors on the 99 concentration of SOAs. Similarly, we used Version 4.7.1 of the CMAO, which is coupled with the 100 gas-phase photochemical mechanism SAPRC99 (1999 Statewide Air Pollutant Research Center 101 (Carter, 2000) and Version 5 of the aerosol module (AERO5) (Byun and Schere, 2006; Foley et al., 102 2010). There are three major formation pathways for SOAs in this version, which is based on the 103 two-product approach. The first pathway is the equilibrium partition of semi-volatile products 104 formed from the oxidation of seven VOC precursors: long-chain alkanes, benzene (BNZ), high-105 yield aromatics (mainly toluene, ARO1), low-yield aromatics (mainly xylene, ARO2), isoprene 106 (ISOP), monoterpene (TRP) and sesquiterpenes (SESQ). The second pathway is the oligomerization 107 of semi-volatile SOAs formed through the first pathway (Kalberer et al., 2004), namely aging 108 process. The third pathway is the formation of SOA via the in-cloud oxidation of glyoxal (GLY) and 109 methylglyoxal (MGLY) (Carlton et al., 2008), both of which represent dicarbonyls in the model. 110 The details of these formation pathways are given in Carlton et al. (2010).

The meteorological fields used to drive CMAQ are obtained from RAMS, which has been described in detail by Cotton et al. (2003). The National Centers for Environmental Prediction reanalysis datasets are served as the initial and lateral boundary meteorological conditions input into RAMS. The boundary conditions used for the RAMS computations include the weekly average sea surface temperature and the monthly measured snow cover. The final modeled results are output through the four-dimensional data assimilation mode using nudging analysis.

The emission sources are derived from several different inventories. Anthropogenic emissions (Li et al., 2017b)—including SO₂, NO_x, CO, black carbon, non-methane VOCs, organic carbon, NH₃ and other particulate matter—are obtained from the monthly emissions inventory of 2012. There are five emission sectors (the power, industrial, residential, transportation and agricultural sectors) in the inventory with a spatial resolution of $0.25^{\circ} \times 0.25^{\circ}$ (see www.meicmodel.org). The Model of 122 Emissions of Gases and Aerosols from Nature (MEGAN) (Guenther et al., 2012) provides the 123 biogenic emissions. The emissions from open biomass burning are derived from the Global Fire 124 Emissions Database, Version 4 (Randerson et al., 2015), whereas the information of monthly lightning NO_x is obtained from the Global Emissions Inventory Activity with a spatial resolution of 125 $1^{\circ} \times 1^{\circ}$ (Benkovitz et al., 1996). The emissions of NO_x from the soil are derived from the Regional 126 Emission inventory in ASia, Version 2.1, with a spatial resolution of $0.25^{\circ} \times 0.25^{\circ}$ (Kurokawa et al., 127 2013). The online dust and sea salt emissions are calculated using the empirical model developed 128 129 by Han et al. (2004) and the scheme of Gong (2003), respectively. The calculation of the model 130 boundary conditions is supported by the Model of Ozone and Related Tracers, Version 4 (MOZART-131 4) (Emmons et al., 2010). The model domain is divided into 105×86 grid cells with the center located at $(35^{\circ} \text{ N}, 110^{\circ} \text{ E})$ 132 133 (Fig. 1). The map projection is rotated polar-stereographic with a horizontal resolution of 64 km.

The vertical simulation region is unequally spaced from the ground surface to ~23 km. There are 25 layers in the σ_z coordinate system in RAMS, with nine layers located in the lowest 2 km to resolve the planetary boundary layer. The vertical dimension of CMAQ is divided into 15 layers, with the lowest seven layers the same as those in RAMS. We focused on the area outlined by the blue box in Fig. 1.

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140 **2.2 Adding the aqueous uptake process**

141 There is a standard reaction probability formulation for reactive uptake of gases by aerosols and 142 clouds in Jacob (2000). In this formulation, the first-order rate constant k for the chemical loss of a 143 gas-phase species to the aerosols or cloud droplets through molecular diffusion and free collision is 144 given by

145 $k = \left(\frac{a}{D_g} + \frac{4}{v\gamma}\right)^{-1} A \tag{1}$

where D_g is the gas-phase molecular diffusion coefficient, *a* is the radius of the aerosol particle or cloud droplet, *v* is the mean molecular speed, *y* is the reaction uptake coefficient when a collision occurs between a gas-phase molecule and the aqueous surface, and *A* is the aqueous particle surface area per unit volume of air.

150 We implement the aqueous uptake of dicarbonyls by cloud droplets following the standard

151 equation (1). In a similar manner to Fu et al. (2008) and Li et al. (2013), the cloud droplet surface 152 area is calculated from the LWC in the cloudy fraction of the model grid by assuming an effective 153 droplet radius of 6 μ m for continental clouds and 10 μ m for maritime clouds. D_g is calculated according to Perry and Green (1999) following Dentener and Crutzen (1993). Similar to previous 154 155 studies (e.g. Lin et al., 2014; Ying et al., 2015; Hu et al., 2017), the uptake of gas-phase dicarbonyls 156 on aqueous aerosols is simply parameterized using the collision limitation ($v\gamma A/4$) of equation (1). The value of $\gamma = 2.9 \times 10^{-3}$ for glyoxal from Liggio et al. (2005) is used in both cloud and aerosol 157 158 processing. While the uptake coefficient adopted for methylglyoxal is from De Haan et al. (2018): the average value of $\gamma = 5.7 \times 10^{-3}$ measured on cloud-processed glycine aerosols is used to account 159 for the uptake by cloud droplets; the average value of $\gamma = 3.0 \times 10^{-3}$ used in aerosol processing is 160 derived from the uptake coefficients measured on deliquesced ammonium sulfate aerosols (3.7×10⁻ 161 ³) and on deliquesced glycine aerosols $(2.3 \times 10^{-3}, \text{ median value})$. 162

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164 **2.3 Observational data**

165 To assess the contribution of dicarbonyl species to the concentration of SOA, we used the observed 166 hourly concentration of SOA in Beijing during the summer and fall of 2014 measured by Xu et al. 167 (2017) with an Aerodyne high-resolution time-of-flight aerosol mass spectrometer. Samples were taken from June 3 to July 11, 2014 (episode 1) and October 14 to November 14, 2014 (episode 2) 168 169 at the Institute of Atmospheric Physics. More detailed information about the data has been reported 170 by Xu et al. (2017). The simulation periods are from May 22 to July 11 and from October 1 to 171 November 14, with the first 12 days as the spin-up time. To evaluate the reasonability in simulating 172 the formation processes of SOAs in the aqueous phase, the corresponding cloud water path (CWP) 173 data measured by the MODerate Resolution Imaging Spectroradiometer (MODIS) was also 174 obtained from the website http://ladsweb.modaps.eosdis.nasa.gov/api/v1/productPage/product=MY 175 DATML2.

Both Liu et al. (2012) and Li et al. (2018) have suggested that the underestimation of glyoxal concentrations in simulations is related to the underpredicted emission of aromatic compounds. Therefore, the biases in the emission of aromatic compounds needs to be evaluated through a comparison of the observed and simulated concentrations of aromatic compounds. The observed data were collected at 14:00 local standard time every Thursday by gas chromatography–mass spectrometer at Beijing, Xinglong and Yucheng (Fig. 1). Sun et al. (2016) and Wu et al. (2016) have
presented the detailed information.

To evaluate the performance of our model, we also compared the simulated PM_{2.5} concentrations and several meteorological factors with the results measured over 12 stations (Fig. 1) during the two episodes. The observed meteorological data were derived from the Meteorological Information Comprehensive Analysis and Process System (MICAPS) dataset, whereas the Chinese National Environmental Monitoring Center provided the measured concentrations of PM_{2.5}.

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189 **3. Results and discussions**

190 **3.1 Model evaluation**

191 Table 1 shows statistical results of the meteorological parameters (i.e. temperature, relative humidity, 192 wind speed and wind direction) and $PM_{2.5}$ concentrations in the two analyzed episodes, where: N is 193 the total number of samples; IOA is the index of agreement, which can synthetically reflect the 194 combination of the modeled value and variable tendency being good or bad (Willmott, 1981); C_{mod} and C_{obs} are the average values of modeled and observed results, respectively; MB and FB are the 195 196 mean and fractional bias, respectively; RMSE and FE are the root-mean-square and fractional error, 197 respectively; and R is the correlation coefficient between the observed and simulated results. The 198 calculations of these statistical parameters can be found in Juda-Rezler et al. (2012). P_{22.5°} and P_{45°} 199 represent the proportions of compared results that the absolute biases between the simulated and 200 measured wind directions are within 22.5° and 45°, respectively (Li et al., 2017a).

201 There are inevitably some biases in the simulated meteorological parameters relative to the 202 observations due to the limited model resolution and system errors. Nevertheless, the model 203 reproduces the magnitude and variation trend of the temperature and relative humidity fairly well 204 (Figs. S1 and S2), with IOAs for temperature of 0.91 and 0.95 in the two episodes and IOAs of 0.91 205 and 0.88 for the relative humidity, comparable with the results of Li et al. (2012) and Wang et al. 206 (2014). The mean biases of temperature $(-0.83^{\circ}C, -0.68^{\circ}C)$ and relative humidity (1.53%, -0.05%)207 are small in the two episodes. The RMSEs of temperature and relative humidity are comparable 208 with the results of Gao et al. (2016) and Zhang et al. (2012), implying a reasonable performance of 209 the simulation. Although the capture of the variable tendency of wind speed (Fig. S3) by the model 210 is rather poor, the small values for the mean bias (-0.40 and -0.22 m/s) and the high IOAs (0.61 and -0.22 m/s)

211 0.69) indicate a better performance than that reported by Feng et al. (2016) and Wang et al. (2014) 212 in simulating the wind speed. The RMSE of wind speed shows that the simulated results achieve 213 the criteria for a good performance ($|RMSE| \le 2 \text{ m/s}$) given by Emery et al. (2001). For the wind 214 direction (Fig. S4), P_{22.5°} and P_{45°} are 32.12 and 54.52% in episode 1 and 34.01 and 57.41% in 215 episode 2, indicating the reasonable simulation of wind direction. Therefore the model provides a 216 reasonable meteorological field for the subsequent simulations.

217 The modeled $PM_{2.5}$ concentrations are generally higher than the observed values in the two 218 episodes over the 12 stations (Fig. S5). Except for the limited resolution of the model, the 219 uncertainties of the emission inventory and the effects of the background transport may also 220 contribute to the overestimation. However, the correlation coefficients of PM_{2.5} are 0.50 and 0.56 in 221 the two episodes, respectively, and the IOAs are 0.64 and 0.70 in the two episodes, respectively, 222 indicating good capture of the variable tendency and magnitude by the model. Both the fractional 223 error and the fractional bias for PM_{2.5} in the two episodes fulfill the performance criteria (FE \leq 224 75%, $|FB| \leq 60\%$) given by Boylan and Russell (2006), implying a good performance in 225 simulating PM_{2.5}. These results show that the model gives a reasonable simulation of the chemical 226 species.

227 As described in Section 2.2, pathway M is related to the cloud water content. Thus the modeled 228 and satellite-observed mean CWP were compared during the two analyzed episodes to evaluate the 229 reasonability of the simulation results. Fig. 2a and 2b show that in episode 1, during one summer 230 period, the highest observed CWP (400-500 g/m²) mainly appears over the Yellow Sea and Bohai 231 Sea and in parts of southeastern and southern China, whereas the observed second highest values 232 (200-300 g/m²) are concentrated in from the Oinghai-Tibetan Plateau to the North China Plain 233 along the Yellow River Basin. The modeled high CWP (100-400 g/m²) appears in the same regions 234 as in the observational dataset. In episode 2, during one fall period (Fig. 2c and 2d), the observed 235 highest (300-500 g/m²) and second highest (200-300 g/m²) CWP values are mainly concentrated in 236 the Hetao area and the regions including the northeastern China and Inner Mongolia, respectively. 237 The modeled values $(50-150 \text{ g/m}^2)$ are also high over these regions.

Overall, there are obvious biases in the numerical values between the observed and simulated CWP. This is a result of the large uncertainties in both the model simulations and the satellite retrievals of clouds, which lead to uncertainties in the simulations of the concentrations of SOAs. However, the mean distribution patterns of the simulated CWP during the two episodes are similar to the observational results, indicating few impacts on the simulated distribution of SOA. Both the simulated and observed CWP in episode 1 are higher than in episode 2, also implying differences in

- the simulation of SOA between the two episodes.
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246 **3.2 Model results and analyses**

As described in Section 2.3 and with reference to previous research, the underestimation of glyoxal concentrations may partly result from the underprediction of the emissions of aromatic compounds. It is therefore necessary to evaluate the emissions of aromatic compounds during the analyzed episodes before designing and implementing the model case studies. For this purpose, comparisons were made between the simulated and observed concentrations of aromatic compounds in a similar manner to the work of Zhang and Ying (2011).

253 Fig. 3 presents the ratio of the observed to predicted (O/P ratio) concentrations of aromatic 254 compounds for the original emissions (Fig. 3a and 3c) and a three-fold increase in the emissions of 255 aromatic compounds (Fig. 3b and 3d) during the two episodes. Fig. 3a shows that there are biases 256 between the observed and simulated concentrations of aromatic compounds in episode 1 at the 257 original emission rates of the aromatic compounds. The O/P ratios are less variable for ARO1 than 258 for ARO2, with both the mean and median O/P values being close to 2. The range of O/P values is 259 large for ARO2 and there is an order of magnitude difference between the lowest and highest ratios. 260 Both the mean and median O/P ratios of ARO2 are about 20. These results show that there are 261 underestimations in the emissions of ARO1 and ARO2. In view of the limitations of the amount of 262 observed data and the nonlinear relationship between emissions and the concentrations (Li et al., 263 2018), the underestimation in the emission rates has been conservatively estimated to be a factor of 264 three. As a result, when the emission rates of aromatic compounds are increased (Fig. 3b), the ratios 265 of ARO1 and ARO2 in episode 1 are less variable, especially ARO2. The mean and median ratios 266 of ARO1 are both around 1, whereas the two ratios for ARO2 decrease from 20 to 6. The results for 267 episode 2 are similar to those in episode 1. Fig. 3c shows that the original O/P ratio of ARO1 varies 268 between 1 and 3.5, with a mean value close to 2, whereas the ratio of ARO2 ranges from 1 to 5, with 269 a mean ratio of about 4, both of which indicate unaccounted emissions of aromatic compounds. Fig. 270 3d shows a clear decrease in the biases of the observed and simulated concentrations of aromatic

271 compounds. The O/P ratios of ARO1 and ARO2 are concentrated and vary between 1 and 2, with a 272 mean ratio close to 1. However, it is difficult to determine whether the factor of 3 is the actual 273 underestimation in the emission of aromatic compounds as a result of its dependence on space and 274 time. For convenience, a factor of 3 is chosen here as a uniform scale with which to assess the 275 unaccounted emissions of aromatic compounds.

276 Three sensitivity simulation case studies are designed based on these results. Case 0 is run with 277 the three default SOA formation pathways included in the standard model. To avoid double-counting 278 the loss of dicarbonyls through the in-cloud oxidation and the following formed SOA, the default 279 aqueous formation pathway is turned off when run case 1 with the pathway M incorporated. 280 According to previous studies (e.g. Stavrakou et al., 2009; Stavrakou et al., 2010; Li et al., 2016), the 281 yields of glyoxal from isoprene are very likely to be underestimated. Li et al. (2018) has run a simple 282 sensitivity case to evaluate the impacts of underestimated glyoxal yields from isoprene on glyoxal 283 simulation. From their results, a better agreement has achieved between the observed and simulated 284 glyoxal after a five-fold increase of glyoxal yields from isoprene. More detailed analyses can be 285 found in Li et al. (2018). Thus, to take the effects of the underestimations of glyoxal concentrations 286 into considerations, case 2 is designed with a three-fold increase in the emissions of aromatic 287 compounds and a five-fold increase in the molar yield of glyoxal from isoprene based on the results 288 of case 1.

Fig. 4 compares the hourly concentrations of the simulated and observed SOAs during the two analyzed episodes; the corresponding statistical parameters are listed in Table 2. The concentrations of SOAs are measured from PM_1 . The observed $PM_1/PM_{2.5}$ ratio of 0.77 (Xu et al., 2015) was used to convert the observed concentrations for comparison with the simulated results for $PM_{2.5}$.

293 In case 0, the SOA concentrations in episode 1 (Fig. 4a) are significantly underestimated by an 294 average factor of 5.7, with the differences being as high as a multiple of ≥ 60 . As a result of the 295 impacts of uncertainties in the meteorological fields and emissions, the variation trend of the 296 concentration of SOA is not well captured by the model (R = 0.21). Similarly, although the variation 297 of SOA is reproduced well in episode 2 (R = 0.83) (Fig. 4b), the concentration of SOA is still 298 underpredicted by an order of magnitude. The biases between the observed and simulated 299 concentrations of SOAs decrease in case 1 when pathway M is added to the model, especially in 300 episode 1. It shows that the mean concentration of SOA in case 1 increases by $3.65 \text{ }\mu\text{g/m}^3$ during

301 episode 1, explaining 34.8% of the unaccounted sources of SOAs. The decreased mean bias and 302 RMSE and the slightly increased IOA also indicate the more reasonable descriptions of the 303 formations of SOAs. In episode 2, the SOA formed through pathway M contributes less to the total 304 concentration of SOA. The mean SOA concentration during episode 2 reaches 2.87 μ g/m³, increased by 1.82 µg/m³. However, the decreased bias statistical parameters (e.g. RMSE and MB) also indicate 305 306 the more realistic description of contributions of dicarbonyls to SOA in this episode. The larger 307 contribution of dicarbonyls from pathway M to the concentration of SOA in episode 1 is attributed 308 to the higher LWC and the larger amount of dicarbonyls produced from biogenic precursors in the 309 summer than in the fall. When the impact of the underestimation of glyoxal on the concentration of 310 SOA is taken into consideration, the concentration of SOA clearly increases in case 2. Compared to 311 case 0, the mean concentration of SOA in episode 1 is significantly improved by a factor of 5.4 and 312 comparable to the observations, whereas the increase in episode 2 is a factor of 6.2. The statistical 313 parameters (e.g. IOA and RMSE) also show the better performance of case 2, indicating a more 314 realistic and reasonable representation of the formation of SOA. Aromatic compounds are not only 315 the precursors of glyoxal, but are also the precursors of SOA in the gas phase. A compositional 316 analysis is therefore required to evaluate the individual contributions from dicarbonyls in case 2. 317 Fig. 5 shows the mean contributions from different sources of SOAs in the three sensitivity case 318 studies during the two analyzed episodes. AAQ, dicarbonyl-derived SOAs, contributes little in case 319 0 during the two episodes. The mean contribution of AAQ to the total concentration of SOAs in 320 episode 1 is 2.39%, larger than in episode 2 (0.89%). As a result of the greater amount of emissions 321 from biogenic sources in summer, SOAs formed from biogenic precursors (AISOP+ATRP+ASQT) 322 contribute more (26.3%) in episode 1 than that in episode 2 (13.5%), whereas the contributions from 323 anthropogenic precursors (AALK+ATOL+AXYL+ABNZ) are comparable between the two 324 episodes. When pathway M is included in the model, the contribution from AAQ to SOA in case 1 325 clearly increases and reaches 57.5% in episode 1 and 63.7% in episode 2. The significant 326 contributions from AAQ in the two episodes indicate the major contributions from dicarbonyls 327 through pathway M to the formations of SOAs in summer and fall. When the impact of the 328 underestimation of glyoxal is taken into consideration in case 2, the mean concentration of AAQ in 329 both episodes are higher than double that of case 1. In episode 1, AAQ $(7.39 \,\mu\text{g/m}^3)$ accounts for 61.0% of the total SOA, exceeding the sum of the contributions from ARO1 (ATOL) and ARO2 330

(AXYL) (15.7%), and indicating the dominant contribution of aqueous-phase process to the concentration of SOA in summer. The increase in AAQ relative to case 0 compensates for about 70.0% of the unaccounted sources of SOAs. In episode 2, AAQ ($4.17 \mu g/m^3$) contributes 64.5% to the total concentration of SOA, which is also higher than the sum of ATOL and AXYL (15.3%), implying the dominant contributions of dicarbonyls to the concentrations of SOAs. The different contribution of AAQ in case 2 during the two episodes can be attributed to the different LWC and different amount of dicarbonyls produced from biogenic precursors.

338 It is clear that the biases between the observed and simulated concentrations of SOAs decrease 339 when the contributions of unaccounted dicarbonyls to the concentrations of SOAs are considered, 340 especially in summer. However, the sources of unaccounted SOAs cannot be explained completely. 341 As a result of uncertainties in the description of known SOA formation processes and missing 342 pathways that are not included in the model-for example, there are many uncertainties in glyoxal 343 simulations (Li et al. (2018)). There are also many uncertainties in incorporating pathway M into 344 the model, such as the effective radius of cloud droplets (the empirical values used instead of the 345 actual values), the reaction uptake coefficient (updated values over disparate land surfaces are 346 discussed in Curry et al. (2018)), and the liquid water content in clouds (the most uncertain 347 parameter shown in Fig. 2). Other pathways for the formations of SOAs, such as the uptake of 348 isoprene epoxides on the surface of aerosols (Lal et al., 2012;Lin et al., 2013), the aging mechanism 349 of semi-volatile primary organic aerosols (Shrivastava et al., 2008) and the oxidation of primary 350 semi- and intermediate VOCs, are not considered in this model. Besides, recent studies (e.g. 351 Galloway et al., 2009; De Haan et al., 2018) have presented that the reactive uptake of glyoxal and 352 methylglyoxal can be reversible, especially methylglyoxal, but we do not consider in this study. 353 To distinguish the contribution of dicarbonyls to the concentration of SOA over China in case 2

from that in case 0, the distributions of dicarbonyl-derived SOAs and their contributions to SOAs
(AAQ/SOA) over China in cases 0 and 2 are analyzed.

Fig. 6(a, b) and 6(c, d) show the mean concentration of AAQ in cases 0 and 2, respectively, during the two episodes. For the base case, in episode 1 (Fig. 6a), the concentration of AAQ over China is $\leq 0.2 \ \mu g/m^3$. The higher concentration of AAQ (0.1–0.2 $\mu g/m^3$) is in the areas between the lower reaches of the Yellow and Yangtze rivers and in the Sichuan Basin, probably due to the higher LWC and a greater number of sources of dicarbonyls over these areas, as discussed in Section 3.1 and by 361 Li et al. (2018). While in episode 2 (Fig. 6b), the concentration of AAO is $\leq 0.1 \, \mu g/m^3$ over the 362 regions for the lower LWC and less sources of dicarbonyls from biogenic precursors. It is clear that 363 the concentration of AAQ is improved in case 2, when pathway M is added and the impact of the 364 underestimation of glyoxal is considered. Overall, the concentration of AAQ is higher in eastern China than in the west during the two episodes. In episode 1 (Fig. 6c), the concentrations of AAQ 365 366 mostly vary from 2 to $15 \,\mu\text{g/m}^3$ over central and eastern China, with the highest value (10–15 $\mu\text{g/m}^3$) concentrated in the areas over the lower reaches of the Yangtze river. The concentration of AAQ in 367 368 western China is $\leq 1 \, \mu g/m^3$, with the lowest value ($\leq 0.1 \, \mu g/m^3$) in the Qinghai-Tibet Plateau, 369 probably because there are few sources of dicarbonyls. In episode 2 (Fig. 6d), the concentrations of 370 AAQ is mostly in a range of $2-10 \ \mu g/m^3$ over central and eastern China, with the highest value (5– 371 $10 \ \mu g/m^3$) concentrated over regions from south of the lower reaches of the Yellow River to the 372 south of Guangzhou Province as well as the Sichuan Basin. The concentration of AAQ in western 373 China is also $\leq 1 \,\mu\text{g/m}^3$, with the lowest value ($\leq 0.1 \,\mu\text{g/m}^3$) in the Qinghai-Tibet Plateau. Outside 374 China, the highest concentration of AAQ (15-20) appears in the northeastern India due to more 375 primary sources of dicarbonyls from the large scale of postharvest paddy residue burnings (Chandra 376 and Sinha, 2016) and the barrier of precursor gases diffusions from Himalayan orogen as well as 377 the low wind speed.

378 Fig. 6(e, f) and 6(g, h) show the spatial distribution of the mean AAQ/SOA in cases 0 and 2 during 379 the two episodes, respectively. Fig. 6e shows that the AAQ fraction over China in episode 1 is \leq 380 10%, except in Yunnan Province and some parts of the South China Sea, where AAQ/SOA reaches 381 10–20%. In episode 2 (Fig. 6f), the mean AAQ/SOA is \leq 10% over the whole regions. When the 382 contributions of dicarbonyls from pathway M and improved glyoxal to SOA are considered, there 383 is a large increase in AAQ/SOA over these regions in two episodes. As shown in Fig. 6g, in episode 384 1, the contribution of AAQ to SOA varies from 10 to 90% throughout China. In central and eastern 385 areas, the fraction of AAQ is generally in the range of 50-70% and up to 70-90% in the coastal 386 regions and offshore from the East China Sea to the South China Sea. The fraction of AAQ in the 387 west is relatively low and usually \leq 50%. However, the contribution of AAQ to SOA is up to 80% 388 in southwestern regions (e.g. Yunnan Province and parts of Sichuan Province). In episode 2 (Fig. 389 6h), the contribution of AAQ to SOA is in the range of 10-80% throughout China. In central and 390 eastern areas, the fraction of AAQ is generally in the range of 50-70% and up 80% in a small portion of northeastern China. The fraction of AAO in the west is also lower and usually \leq 50%. However,

the contribution of AAQ to SOA is up to 60-70% in a small part of Sinkiang Province.

393 4. Conclusions

394 The RAMS-CMAQ modeling system was used to assess the contributions of dicarbonyls from the 395 aqueous uptake process and unaccounted sources of glyoxal to the concentrations of SOAs during 396 the two episodes from June 3 to July 11, 2014 and October 14 to November 14, 2014. Comparisons 397 between the observed and simulated concentrations of SOAs from three sensitivity groups showed 398 different improvements in the SOA simulations with the inclusion of pathway M and consideration 399 of the underestimation of glyoxal in the two episodes. Due to the high LWC and large amount of 400 dicarbonyls produced from biogenic precursors in summer, the contributions of dicarbonyls were 401 greater in episode 1 than that in episode 2. When pathway M was added in case 1, the modeled mean 402 concentration of SOA in episode 1 increased by $3.65 \text{ }\mu\text{g/m}^3$, explaining about 34.8% of the 403 unaccounted SOA sources, while there was a $1.82 \,\mu g/m^3$ increase in the mean result during episode 404 2. When the impacts of glyoxal underestimation were taken into consideration in case 2, the modeled 405 mean AAQ in episode 1 was improved to 7.39 $\mu g/m^3$ and contributed 61.0% of the total 406 concentration of SOA. The increase due to AAQ relative to case 0 is equivalent to 70.0% of the 407 unaccounted source of SOA. Whereas the mean concentration of AAQ in episode 2 was $4.17 \,\mu\text{g/m}^3$ 408 and accounted for 64.5% of total concentration of SOA. Although the contributions of dicarbonyls 409 to SOAs are different in the two episodes, the simulated SOA results are both improved and close 410 to the observations, indicating a more realistic description of aqueous formation of SOA.

411 The mean AAQ in case 2 during the two episodes was clearly improved over China relative to 412 case 0 and was generally higher in the east than in the west. In episode 1, the highest value (10–15 413 $\mu g/m^3$) was seen in the areas around the lower reaches of the Yellow River. While the highest value 414 $(5-10 \ \mu g/m^3)$ in episode 2 was concentrated over regions from south of the lower reaches of the 415 Yellow River to the south of Guangzhou Province as well as the Sichuan Basin. As a result, the 416 contribution of AAQ to the concentration of SOA in two episodes was also improved in case 2. In 417 episode 1, the fraction varied from 10 to 90% throughout China, with the highest contribution (70-418 90%) in the coastal regions and offshore along the East China Sea to the South China Sea in addition 419 to the southwestern regions (e.g. Yunnan Province and parts of Sichuan Province). While in episode 420 2, the contribution of AAQ to SOA was in the range of 10–80% throughout China, with the highest 421 fraction (up to 80%) seen in a small portion of northeastern China.

It is clear that the contributions of dicarbonyls from pathway M and the unaccounted sources of glyoxal make a significant contribution to the concentration of SOA, especially in summer. However, there are still many uncertainties in the depictions of aqueous uptake processes and sources of glyoxal. More work is needed on the parameters used in pathway M (e.g. the effective radius of cloud droplets and the reaction uptake coefficient), and to improve the accuracy of the modeled liquid water content in clouds.

428

429 *Author contribution.* In this study, J. Li designed the sensitivity experiments, developed the model code, and 430 performed the corresponding simulations. M. Zhang co-designed the experiments and provided valuable advices 431 about the model operations. Y. Sun carried out the measurements of SOA and provided the corresponding data for 432 evaluating the modeled results. G. Tang and F. Wu carried out the measurements of aromatic compound 433 concentrations and provided the corresponding data for evaluating the simulated results. Y. Xu provided valuable 434 advice on model result analysis.

435

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439 dicarbonyls pathway into the model.

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719 Table 1. Statistics for the meteorological variables and PM_{2.5} between the simulated and observed data during the

720	two episodes.
120	two episodes.

Table 1-1								
Variables	Episode	N	$C_{ m mod}$	$C_{\rm obs}$	MB	RMSE	R	IOA
Temperature	1	10770	23.76	24.59	-0.83	2.58	0.86	0.91
(°C)	2	8391	12.68	13.36	-0.68	3.13	0.92	0.95
Relative humidity	1	10770	69.42	67.90	1.52	13.68	0.83	0.91
(%)	2	8389	62.81	62.86	-0.05	14.73	0.77	0.88
Wind speed	1	10770	1.78	2.18	-0.40	1.64	0.35	0.61
(m/s)	2	8391	2.05	2.27	-0.22	1.75	0.50	0.69
Table 1-2								
Variables	Episode	N	$C_{ m mod}$	$C_{\rm obs}$	FB (%)	FE (%)	R	IOA
PM _{2.5}	1	8091	64.04	47.90	13.35	64.61	0.50	0.64
$(\mu g/m^3)$	2	7649	86.25	61.67	36.00	62.27	0.56	0.70
Table 1-3								
Variables	Episode	Ν		P _{22.5°} (%)		$P_{45^{\circ}}$ (%)		
Wind direction	1	10770		32.12		54.52		
(°)	2	8391		34.01		57.41		

N is the total number of samples; C_{mod} and C_{obs} are the average value of modeled and observed results, respectively;

MB and FB are the mean and fractional bias, respectively; RMSE and FE are the root-mean-square and fractional error, respectively; IOA is the index of agreement; R is the correlation coefficient between the observed and simulated results; P_{22.5°} and P_{45°} represent the proportions of compared results that the absolute biases between the simulated and measured wind directions are within 22.5° and 45°, respectively.

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Table 2. Performance statistics of the modeled and observed SOA concentrations ($\mu g/m^3$).

Case	Episode	N	C_{model}	$C_{\rm obs}$	MB	RMSE	R	IOA
0	1	822	2.21	12.69	-10.48	13.48	0.21	0.46
	2	737	1.05	17.69	-16.64	23.52	0.83	0.45
1	1	822	5.86	12.69	-6.83	10.86	0.25	0.49
	2	737	2.87	17.69	-14.82	21.21	0.84	0.50
2	1	822	12.03	12.69	-0.66	10.19	0.21	0.52
	2	737	6.53	17.69	-11.16	16.56	0.83	0.64

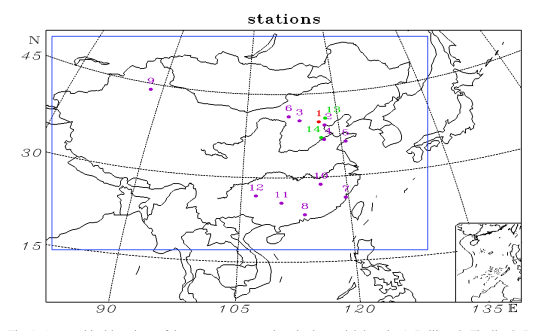
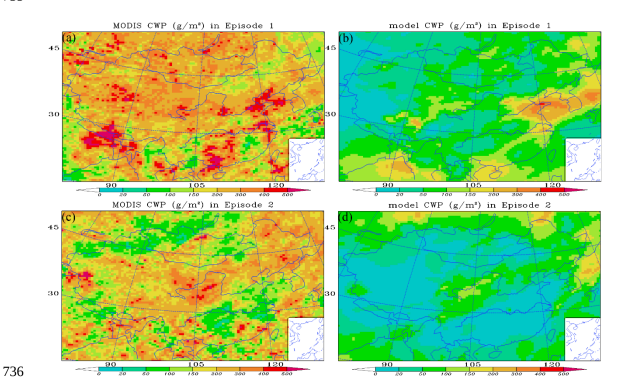


Fig. 1. Geographical locations of the measurement stations in the model domain. 1: Beijing; 2: Tianjin; 3: Datong;
4: Jinan; 5: Qingdao; 6: Hohhot; 7: Fuzhou; 8: Guangzhou; 9: Urumqi; 10: Nanchang; 11: Guilin; 12: Guiyang; 13:
Xinglong; 14: Yucheng. The different colors denote which variables are compared at each station: purple:
meteorological parameters; green: concentrations of aromatics; red: both meteorological parameters and aromatic
compound concentrations.





737 Fig. 2. Observed and simulated distributions of the mean cloud water path (CWP) during the two episodes (left-hand

738 panels: MODIS observation; right-hand panels: results of simulations).

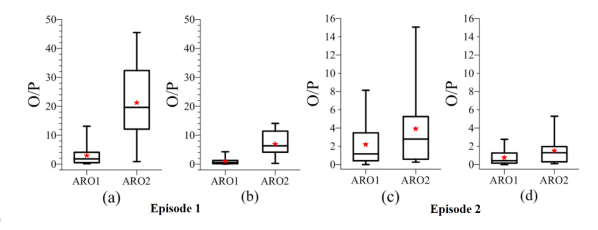




Fig. 3. Box-whisker plot of the observed to predicted (O/P) ratios of aromatic compounds during the two analyzed
episodes. The observed data were measured at 14:00 LST every Thursday by gas chromatography-mass spectrometer
at three sites (Beijing, Xinglong and Yucheng). (a, c): Original emissions of aromatic compounds; and (b, d): a threefold increase in the emissions of aromatic compounds. The red stars show the mean O/P ratios.

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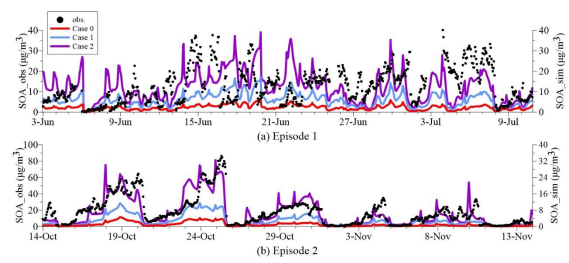
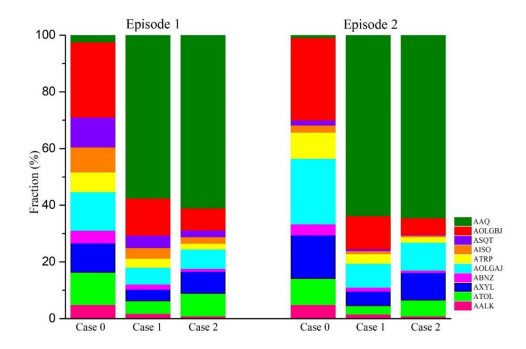


Fig. 4. Hourly concentrations of the observed and simulated near-surface SOA concentrations in episode 1 (a) and 2
(b). Case 0 is base run; case 1 is run with the incorporation of the aqueous uptake of dicarbonyls (excluding the
default in-cloud dicarbonyl oxidations); and case 2 is run with the glyoxal underestimation taken into consideration
based on case 1.



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Fig. 5. Mean contributions from different sources of SOAs in the three sensitivity case studies during the two analyzed episodes. The compositions represent the SOA formed from long-chain alkanes (AALK), high-yield aromatic compounds (ATOL), low-yield aromatic compounds (AXYL), benzene (ABNZ), monoterpenes (ATRP), isoprene (AISO) and dicarbonyls (AAQ), as well as aged anthropogenic (AOLGAJ) and biogenic SOA (AOLGBJ). Case 0 is the base run; case 1 is run with the incorporation of the aqueous uptake of dicarbonyls (excluding the default in-cloud dicarbonyl oxidations); and case 2 is based on case 1, but taking into consideration the underestimation of glyoxal.

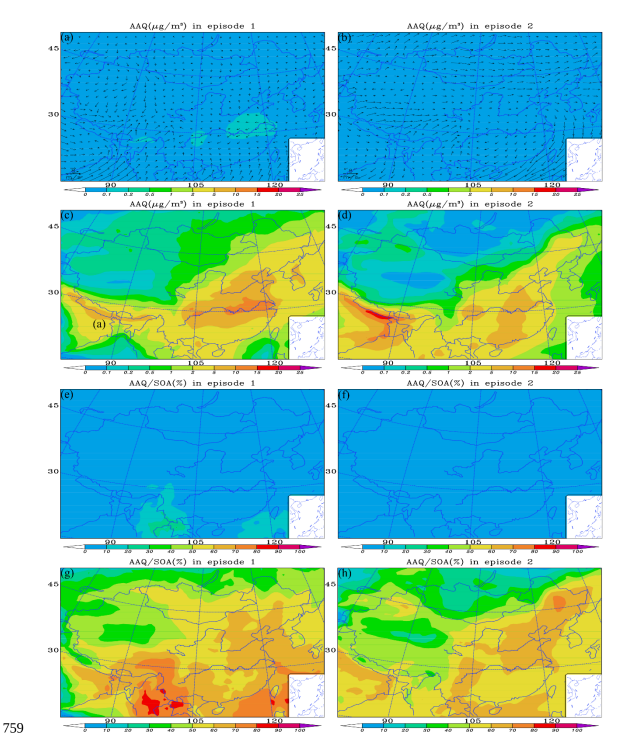


Fig. 6. Modeled distributions of the mean (a, b) wind field with SOA formed from dicarbonyls (AAQ) in case 0, (c,
d) AAQ in case 2, (e, f) AAQ/SOA in case 0, (g, h) AAQ/SOA in case 2, over the regions during two episodes. Case
0 is the base run; case 2 is run taking into consideration the incorporation of the aqueous uptake of dicarbonyls
(excluding the default in-cloud dicarbonyl oxidations) and the underestimation of glyoxal.